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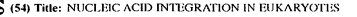
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(57) Abstract: The invention relate to the field of molecular biology and cell biology. It particularly relates to methods to direct integration of a nucleic acid of interest towards homologous recombination and uses thereof. The present invention discloses factors involved in integration of a nucleic acid by illegitimate recombination which provides a method to direct integration of a nucleic acid of interest to a pre-determined site, whereby said nucleic acid has homology at or around the said pre-determined site, in a eukaryote with a preference for non-homologous recombination comprising steering an integration pathway towards homologous recombination. Furthermore, the invention provides a method to direct integration of a nucleic acid of interest to a sub-telomeric and/or telomeric region in a eukaryote with a preference for non-homologous recombination.

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Title: Nucleic acid integration in eukaryotes

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The invention relates to the field of molecular biology and cell biology. It particularly relates to methods to direct integration towards homologous recombination and uses thereof. Several methods are known to transfer nucleic acids to, in particular, eukaryotic cells. In some methods the nucleic acid of interest is transferred to the cytoplasm of the cell, in some the nucleic acid of interest is integrated into the genome of the host. Many different vehicles for transfer of the nucleic acid are known. For different kinds of cells, different systems can be used, although many systems are more widely applicable than just a certain kind of cells. In plants, e.g., a system based on Agrobacterium tumefaciens is often applied. This system is one of the systems that is used in a method according to the invention.

The soil bacterium Agrobacterium tumefaciens is able to transfer part of its tumor-inducing (Ti) plasmid, the transferred (T-) DNA, to plant cells. This results in crown gall tumor formation on plants due to expression of onc-genes, which are present on the T-DNA. Virulence (vir) genes, located elsewhere on the Ti-plasmid, mediate T-DNA transfer to the plant cell. Some Vir proteins accompany the T-DNA during its transfer to the plant cell to protect the T-DNA and to mediate its transfer to the plant nucleus. Once in the plant nucleus, the T-DNA is integrated at a random position into the plant genome (reviewed by (Hooykaas and Beijersbergen, 1994), (Hansen and Chilton 1999). Removal of the onc-genes from the T-DNA does not inactivate T-DNA transfer. T-DNA, disarmed in this way, is now the preferred vector for the genetic modification of plants.

Although much is known about the transformation process, not much is known about the process by which the T-DNA is integrated into the plant genome. It is likely that plant enzymes mediate this step of the transformation process (Bundock at al. 1995). The integration pattern of T-DNA in transformed plants has been extensively studied (Matsumoto at al. 1990) (Gheysen et al 1991) (Meyerhofer et al 1991). The results indicated that T-

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DNA integrates via illegitimate recombination (IR) (also called non-homologous recombination, both terms may be used interchangeable herein), a process which can join two DNA molecules that share little or no homology (here the T-DNA and plant target DNA). Even T-DNA molecules in which a large segment of homologous plant DNA was present, integrated mainly by IR and only with very low frequency (1:104–105) by homologous recombination (HR) (Offringa et al. 1990).

Recently, it was shown that Agrobacterium is not only able to transfer its T-DNA to plant cells, but also to other eukaryotes, including the yeast S.cerevisiae (Bundock et al. 1995) and a wide variety of filamentous fungi (de Groot et al. 1998). In S.cerevisiae, T-DNA carrying homology with the yeast genome integrates via HR (Bundock et al. 1995). However, T-DNA lacking any homology with the S.cerevisiae genome becomes integrated at random positions in the genome by the same IR process as is used in plants (Bundock and Hooykaas 1996). Apparently, eukaryotic cells have at least two separate pathways (one via homologous and one via non-homologous recombination) through which nucleic acids (in particular of course DNA), can be integrated into the host genome. The site of integration into a host cell genome is important with respect to the likelihood of transcription and/or expression of the integrated nucleic acid. The present invention provides methods and means to direct nucleic acid integration to a pre-determined site through steering integration towards the homologous recombination pathway. The present invention arrives at such steering either by enhancing the HR pathway, or by inhibiting (meaning reducing) the IR pathway.

Host factors involved in the integration of nucleic acid by IR have so far not been identified. The present invention discloses such factors which enables the design of methods for their (temporary) inhibition, so that integration of nucleic acid by IR is prevented or more preferably completely inhibited, shifting the integration process towards HR and facilitating the isolation of a host cell with nucleic acid integrated by HR at a predetermined site. This is

extremely important, since there is no method available yet for easy and precise genetic modification of a host cell using HR (gene targeting). Of course the actual site of integration is then determined by homology of the nucleic acid of interest with said site.

In a first embodiment the invention provides a method to direct nucleic 5 acid integration of a nucleic acid of interest to a pre-determined site, whereby said nucleic acid has homology at or around the said pre-determined site, in a eukaryote with a preference for non-homologous recombination comprising steering an integration pathway towards homologous recombination. Preferably, such a method comprises at least the steps of introducing said 10 nucleic acid of interest to a cell of said eukaryote, for example by the process of transformation or electroporation, and integration of said nucleic acid in the genetic material of said cell. Integration is a complex process wherein a nucleic acid sequence becomes part of the genetic material of a host cell. One step in the process of nucleic acid integration is recombination; via recombination 15 nucleic acid sequences are exchanged or inserted and the introduced nucleic acid becomes part of the genetic material of a host cell. In principle two different ways of recombination are possible: homologous and illegitimate or non-homologous recombination. Most (higher) eukaryotes do not or at least not significantly practise homologous recombination although the essential 20 proteins to accomplish such a process are available. One reason for this phenomenon is that frequent use of homologous recombination in (higher) eukaryotes could lead to undesirable chromosomal rearrangements due to the presence of repetitive nucleic acid sequences. To accomplish homologous recombination via a method according to the invention, it is important to 25 provide a nucleic acid which has homology with a pre-determined site. It is clear to a person skilled in the art that the percentage of homology and the length of (a) homologous region(s) play(s) an important role in the process of homologous recombination. The percentage of homology is preferably close to

100%. A person skilled in the art is aware of the fact that lower percentage of

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homology are also used in the field of homologous recombination, but dependent on, for example, the regions of homology and their overall distribution, can lead to a lower efficiency of homologous recombination but are still useful and therefore included in the present invention. Furthermore, the length of a (nearly) homologous region is approximately 3 kb which is sufficient to direct homologous recombination. At least one homologous region is necessary for recombination but more preferably 2 homologous regions flanking the nucleic acid of interest are used for targeted integration. The researcher skilled in the art knows how to select the proper percentage of homology, the length of homology and the amount of homologous regions. By providing such a homology a nucleic acid is integrated at every desired position within the genetic material of a host cell. It is clear to a person skilled in the art that the invention as disclosed herein is used to direct any nucleic acid (preferably DNA) to any pre-determined site as long as the length of homology and percentage of homology are high enough to provide/enable homologous recombination. A pre-determined site is herein defined as a site within the genetic material contained by a host cell to which a nucleic acid with homology to this same site is integrated with a method according to the invention. It was not until the present invention that a nucleic acid is integrated at every desired position and therefore a method according to the invention is applied, for example, to affect the gene function in various ways, not only for complete inactivation but also to mediate changes in the expression level or in the regulation of expression, changes in protein activity or the subcellular targeting of an encoded protein. Complete inactivation, which can usually not be accomplished by existing methods such as antisense technology or RNAi technology (Zrenner et al, 1993), is useful for instance for the inactivation of genes controlling undesired side branches of metabolic pathways, for instance to increase the quality of bulk products such as starch. or to increase the production of specific secondary metabolites or to inhibit formation of unwanted metabolites. A method according to the invention is

also used to inactivate genes controlling senescence in fruits and flowers or that determine flower pigments. Replacement of existing regulatory sequences by alternative regulatory sequences is used to alter expression of in situ modified genes to meet requirements, (e.g. expression in response to particular physical conditions such as light, drought or pathogen infection, or in response 5 to chemical inducers), or depending on the developmental state (e.g. in a storage organ, or in fruits or seeds) or on tissue or cell types. Also a method according to the invention is used to effectuate predictable expression of transgenes encoding novel products, for example by replacing existing coding sequences of genes giving a desired expression profile by those for a desired 10 novel product. For example to produce proteins of medicinal or industrial value in the seeds of plants the coding sequence of a strongly expressed storage protein may be replaced by that of the desired protein. As another example existing coding sequences are modified so that the protein encoded has optimized characteristics for instance to make a plant herbicide tolerant, to 15 produce storage proteins with enhanced nutritional value, or to target a protein of interest to an organelle or to secrete it to the extracellular space. As yet another example eukaryotic cells (including yeast, fungus, plant, mammalian cells or (non-human) animal cells) are provided with a gene encoding a protein of interest integrated into the genome at a site which 20 ensures high expression levels. As another example the nucleic acid of interest can be part of a gene delivery vehicle to deliver a gene of interest to a eukaryotic cell in vitro or in vivo. In this way a defect p53 can be replaced by an intact p53. In this way a tumoricidal gene is delivered to a pre-determined site present only in e.g. proliferating cells, or present only in tumor cells, e.g. 25 to the site where a tumor antigen is expressed from. Gene delivery vehicles are well known in the art and include adenoviral vehicles, retroviral vehicles, nonviral vehicles such as liposomes, etc. As another example the invention is used to produce transgenic organisms. Knock-out transgenics are already produced by homologous recombination methods. The present invention improves the 30

efficiency of such methods. Also transgenics with desired properties are made. It is clear to a person skilled in the art that transgenics can for example be made by the use of Agrobacterium as gene delivery vehicle for plant (Vergunst et al., 1998), yeast (Bundock et al., 1995), fungus (de Groot et al., 1998) or 5 animal (Kunik et al., 2001) or by direct DNA delivery methods exemplified by but not restricted to electroporation for yeast (Gietz & Woods, 2001), plant (D'Halluin et al., 1992; Lin et al., 1997), fungus (Ozeki et al, 1994) and animal (Templeton et al., 1997), LiCl treatment for yeast (Schiestl et al., 1993), microinjection for plant (Schnorf et al., 1991) and animal (Capecchi, 1980) and 10 "DNA whiskers" for plant (Kaeppler et al., 1990; Dunwell, 1999) or particle bombardment for plants and animals (Klein et al., 1992). It is furthermore clear that transgenic plants can be obtained via selective regeneration of transformed plant cells into a complete fertile plant (Vergunst et al., 1998) or via non-regenerative approaches by transforming germ line cells, exemplified 15 by but not restricted to dipping Arabidopsis flowers into an Agrobacterium suspension (Bechtold et al., 1993). It is also clear that transgenic animals can be obtained by transforming embryonic stem cells with one of the DNA delivery methods mentioned above (Hooper, 1992).

In another embodiment the invention provides a method to direct

20 nucleic acid integration to a pre-determined site, whereby said nucleic acid has
homology at or around the said pre-determined site, in a eukaryote with a
preference for non-homologous recombination comprising steering an
integration pathway towards homologous recombination by providing a
mutant of a component involved in non-homologous recombination. Methods to

25 identify components involved in non-homologous recombination are outlined in
the present description wherein S.cerevisiae was used as a model system. To
this end several yeast derivatives defective for genes known to be involved in
various recombination processes were constructed and the effect of the
mutations on T-DNA integration by either HR or IR was tested. The results as
30 disclosed herein show that the proteins encoded by YKU70, RAD50, MRE11,

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XRS2, LIG4 and SIR4 play an essential role in DNA integration by IR but not by HR. WO 00/12716 describes a maize Ku70 orthologue and suggests that "Control of homologous recombination or non-homologous end joining by modulating Ku provides the means to modulate the efficiency which 5 heterologous nucleic acids are incorporated into the genomes of a target plant cell." WO 00/68404 describes a maize Rad50 orthologue and suggest an analogous control for Rad50. Both patent applications do however not disclose, in contrast to the present patent application, that by preventing or more preferably completely inhibiting non-homologous recombination, for example 10 by providing a mutant of a component involved in non-homologous recombination or by inhibiting such a component, that the integration pathway is steered towards homologous recombination. It is clear to a person skilled in the art that different mutants of a component involved in nonhomologous recombination exist. Examples are deletion mutants, knock-out 15 (for example via insertion) mutants or point mutants. Irrespective of the kind of mutant it is important that a component involved in non-homologous recombination is no longer capable or at least significantly less capable to perform it's function in the process of non-homologous recombination. As disclosed herein disruption of YKU70, RAD50, MRE11, XRS2, LIG4 and SIR4 20 did not affect the frequency of DNA integration by HR, showing that these genes are not involved in DNA integration by HR, but only in DNA integration by IR. More over, in the wild-type yeast strain 85% of the integration events occurred by HR (37% by replacement and 63% by insertion) and 15% by IR. In contrast, integration occurred only by HR in yeast strains lacking ku70 or lig4. 25 In rad50 and xrs2 mutant strains the T-DNA preferentially integrated by HR (92%) and 93% of these T-DNAs integrated by replacement and only 7% by insertion. Thus, the absence of a functional rad50 or xrs2 gene leads to a significantly increased frequency of replacement reactions.

In another embodiment the invention provides a method to direct integration of a nucleic acid of interest to a sub-telomeric and/or telomeric

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region in a eukaryote with a preference for non-homologous recombination by providing a mutant of a component involved in non-homologous recombination. A telomeric region is typically defined as a region containing repetitive sequences which is located at the end of a chromosome. A sub-telomeric region is typically defined as a region flanking the telomeric region. As an example it is disclosed herein that in yeast strains carrying disruptions of RAD50, MRE11 or XRS2 the distribution of integrated DNA copies is altered when compared to wildtype. DNA becomes preferentially integrated in telomeres or subtelomeric regions in the rad50, mre11 and xrs2 mutants. A great advantage of integration of DNA copies in telomeres or subtelomeric regions instead of integration elsewhere in the genomic material is that there is no danger for host genes being mutated or inactivated by a DNA insertion. When in plants deficient for RAD50, MRE11 or XRS2 DNA copies also integrate into telomeres or subtelomeric regions, such plants are used for (sub)telomeric targeting of T-DNA in transformation experiments to prevent additional insertion mutations from random T-DNA integration into the plant genome.

In yet another embodiment the invention provides a method to direct nucleic acid integration to a pre-determined site, whereby said nucleic acid has homology at or around the said pre-determined site, in a eukaryote with a preference for non-homologous recombination comprising steering an integration pathway towards homologous recombination by partially or more preferably completely inhibiting a component involved in non-homologous recombination. Partial or complete inhibition of a component involved in non-homologous recombination is obtained by different methods, for example by an antibody directed against such a component or a chemical inhibitor or a protein inhibitor or peptide inhibitor or an antisense molecule or an RNAi molecule. Irrespective of the kind of (partial or more preferably complete) inhibition it is important that a component involved in non-homologous recombination is no longer capable or at least significantly less capable to perform it's function in the process of non-homologous recombination. In yet

another embodiment the invention provides a method to direct integration of a nucleic acid of interest to a sub-telomeric and/or telomeric region in a eukaryote with a preference for non-homologous recombination by partially or more preferably completely inhibiting a component involved in non-homologous recombination. Preferably, said component involved in non-homologous recombination is rad50, mre11 or xrs2.

In a preferred embodiment the invention provides a method to direct nucleic acid integration to a pre-determined site or to a sub-telomeric and/or telomeric region by providing a mutant of a component involved in nonhomologous recombination or by partially or more preferably completely 10 inhibiting a component involved in non-homologous recombination wherein said component comprises ku70, rad50, mre11, xrs2, lig4, sir4 or others such as ku80 (Tacciole et al., 1994; Milne et al., 1996), lif1 (Teo and Jackson, 2000; XRCC4 in human, see figure 6; Junop et al., 2000) and nej1, (Kegel et al., 2001; Valencia et al., 2001). Components involved in non-homologous 15 recombination are identified as outlined in the present description. The nomenclature for genes as used above is specific for yeast. Because the nomenclature of genes differs between organisms a functional equivalent or a functional homologue (for example NBS1, a human xrs2 equivalent (Paull and Gellert, 1999) and see for example figure 2 to 5) and/or a functional fragment 20 thereof, all defined herein as being capable of performing (in function, not in amount) at least one function of the yeast genes ku70, rad50, mre11, xrs2, lig4, sir4, ku80, lif1 or nej1 are also included in the present invention. A mutant of a component directly associating with a component involved in nonhomologous recombination or (partial or complete) inhibition of a component 25 directly associating with a component involved in non-homologous recombination is also part of this invention. Such a component directly associating with a component involved in non-homologous recombination is, for example, identified in a yeast two hybrid screening. An example of a component directly associating with a component involved in non-homologous 30

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recombination is KU80, which forms a complex with KU70. In a more preferred embodiment the invention provides a method to direct nucleic acid integration in yeast, fungus, plant or (non-human) animal (cells).

In another embodiment the invention provides a method to direct nucleic acid integration to a pre-determined site, whereby said nucleic acid has homology at or around the said pre-determined site, in a eukaryote with apreference for non-homologous recombination comprising steering an integration pathway towards homologous recombination by transiently (partially or more preferably completely) inhibiting integration via nonhomologous recombination. In yet another embodiment the invention provides a method to direct integration of a nucleic acid of interest to a subtelomeric and/or telomeric region in a eukaryote with a preference for non-homologous recombination by transiently (partially or more preferably completely) inhibiting integration via non-homologous recombination. In a more preferred embodiment, such a method is used for yeast, plant, fungus or (non-human) animal and the transient (partial or more preferably complete) inhibition is provided by a (preferably stably) inserted and expressed chimeric transgene that encodes a peptide inhibitory to one, some or all non-homologous recombination (NHR) enzymes fused to a nuclear localisation signal (Hanover, 1992; Raikhel, 1992) and the steroid-binding domain of a steroid receptor (Picard et al., 1988). The chimeric transgene is constructed in such a way, using either heterologous or non-heterologous promoter sequences and other expression signals, that it provides (stable) expression in the target cells or tissue for transformation. In the absence of the steroid hormone, the steroidbinding domain binds to chaperone proteins, and thereby the fusion protein is retained in the cytoplasm. Upon treatment with the steroid hormone, the chaperones are released from the steroid-binding domain and the inhibitory peptide will enter the nucleus where it will interact with and inhibit the action of NHR enzymes. An example of an inhibitory peptide is a KU80 fragment that WO 02/052026 PCT/NL01/00936

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imparts radiosensitivity to Chinese hamster ovary cells (Marangoni et al., 2000).

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In a more preferred embodiment such a method is used for yeast, plant, fungus or a (non-human) animal and the transient (partial or more preferable complete) inhibition is provided by an Agrobacterium Vir-fusion protein capable of (partially or more preferably completely) inhibiting a component involved in non-homologous recombination or capable of (partially or more preferably completely) inhibiting a functional equivalent or homologue thereof or capable of (partially or more preferably completely) inhibiting a component directly associating with a component involved in non-homologous recombination. In an even more preferred embodiment such an Agrobacterium Vir fusion protein comprises VirF or VirE2. It was shown that the Agrobacterium VirF and VirE2 proteins are directly transferred from Agrobacterium to plant cells during plant transformation (Vergunst et al. 2000). To, for example, accomplish T-DNA integration by HR in plants, VirF fusion proteins containing for example a peptide inhibitor of IR in plant cells are introduced concomitantly with the targeting T-DNA. It has been reported that the C-terminal part (approximately 40 amino acids) of VirF or VirE2 is sufficient to accomplish transfer of T-DNA. A functional fragment and/or a functional equivalent of VirF or VirE is therefore also included in the present invention. Preferably, said nucleic acid of interest is delivered to a cell of said eukaryote by Agrobacterium.

In an even more preferred embodiment a component involved in non-homologous recombination comprises ku70, rad50, mre11, xrs2, lig4, sir4, ku80, lif1 or nej1 or functional equivalents or homologous thereof or associating components. The nomenclature for genes as used above is specific for yeast. Because the nomenclature of genes differs between organisms a functional equivalent or a functional homologue (see for example figure 2 to 5) and/or a functional fragment thereof, all defined herein as being capable of performing (in function, not in amount) at least one function of the yeast genes

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ku70, rad50, mre11, xrs2, lig4, sir4, ku80, lif1 or nej1are also included in the present invention. By transiently (partially or more preferably completely) inhibiting a component involved in non-homologous recombination a nucleic acid is integrated at any desired position without permanently modifying a component involved in non-homologous recombination and preventing unwanted side effects caused by the permanent presence of such a modified component involved in non-homologous recombination.

Methods according to the present invention, as extensively but not limiting discussed above, are used in a wide variety of applications. One embodiment of the present invention is the replacement of an active gene by an inactive gene according to a method of the invention. Complete inactivation, which can usually not be accomplished by existing methods such as antisense technology or RNAi technology, is useful for instance for the inactivation of genes controlling undesired side branches of metabolic pathways, for instance to increase the quality of bulk products such as starch, or to increase the production of specific secondary metabolites or to inhibit formation of unwanted metabolites. Also to inactivate genes controlling senescence in fruits and flowers or that determine flower pigments. Another embodiment of the present invention is the replacement of an inactive gene by an active gene. One example is the replacement of a defect p53 by an intact p53. Many tumors acquire a mutation in p53 during their development which renders it inactive and often correlates with a poor response to cancer therapy. By replacing the defect p53 by an intact p53, for example via gene therapy, conventional anti cancer therapy have better changes of succeeding. In yet another embodiment of the invention a therapeutic proteinaceous substance is integrated via a method of the invention. In this way a tumoricidal gene ise delivered to a pre-determined site present only in e.g. proliferating cells, or present only in tumor cells, e.g. to the site where a tumor antigen is expressed from. In yet another embodiment the invention provides a method to introduce a substance conferring resistance for an antibiotic substance to a cell according WO 02/052026 PCT/NL01/00936

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to a method of the invention. Also a method according to the invention is used to confer a desired property to an eukaryotic cell. In a preferred embodiment a gene delivery vehicle is used to deliver a desired nucleic acid to a predetermined site. Gene delivery vehicles are well known in the art and include adenoviral vehicles, retroviral vehicles, non-viral vehicles such as liposomes, etc.. In this way, a for example, tumoricidal gene can be delivered to a predetermined site present only in e.g. proliferating cells, or present only in tumor cells, e.g. to the site where a tumor antigen is expressed from.

Furthermore a method according to the invention is used to improve gene targeting efficiency. Such a method is used to improve for example the gene targeting efficiency in plants. In plants transgenes integrate randomly into the genome by IR (Mayerhof et al. 1991), (Gheysen et al. 1991). The efficiency of integration by HR is very low, even when large stretches of homology between the transgene and the genomic target site are present (Offringa et al. 1990). Therefore, the efficiency of gene targeting using HR is very low in plants. The results that are disclosed herein show how to improve the gene targeting efficiency in plants. From the fact that T-DNA integration by IR is strongly reduced in KU70, RAD50, MRE11, XRS2, LIG4 and SIR4 deficient yeast strains and T-DNA integration by HR is not affected in such strains, T-DNA integration by HR is more easily obtained in plants, deficient for either of these genes. Recently, we have cloned a KU70 homologue of Arabidopsis thaliana (see figure 2, Bundock 2000, unpublished data). RAD50, MRE11 and LIG4 homologues have already been found in A.thaliana (GenBank accession numbers AF168748, AJ243822 and AF233527, respectively, see also figure 3, 4 and 5 (Hartung and Puchta 1999). Currently, screenings are being performed to find plants carrying a T-DNA inserted in AtMRE11, AtKU70 or AtLIG4. These knockout plants are used to test whether T-DNA integration by IR is reduced and integration by HR is essentially unaffected, thereby facilitating the detection of T-DNA integration by HR.

Furthermore, the invention provides a method to direct integration of a nucleic acid of interest to a pre-determined site, whereby said nucleic acid has homology at or around the said pre-determined site, in a eukaryote with a preference for non-homologous recombination, comprising steering an integration pathway towards homologous recombination, wherein said nucleic acid sequence of interest is essentially replacing a sequence within said eukaryote. As disclosed herein within the experimental part, in the wild-type yeast strain 85% of the integration events occurred by HR (37% by replacement and 63% by insertion) and 15% by IR. In contrast, integration occurred only by HR in yeast strains lacking ku70 or lig4. In rad50 and xrs2 mutant strains the T-DNA preferentially integrated by HR (92%) and 93% of these T-DNAs integrated by replacement and only 7% by insertion. Thus, the absence of a functional rad50 or xrs2 gene leads to a significantly increased frequency of the desired replacement reactions.

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The invention will be explained in more detail in the following description, which is not limiting the invention.

EXPERIMENTAL PART

Yeast strains.

The yeast strains that were used are listed in Table 1. Yeast mutants isogenic to the haploid YPH250 strain were constructed using the one-step 5 disruption method (Rothstein 1991). A 1987 bp fragment from the YKU70 locus was amplified by PCR using the primers hdf1p1 5'-GGGATTGCTTTAAGGTAG-3' and hdf1p2 5'-CAAATACCCTACCCTACC-3'. The PCR product was cloned into pT7Blue (Novagen) to obtain 10 pT7Blue YKU70. A 1177 bp EcoRV/HindIII fragment from the YKU70 ORF was replaced by a 2033 bp HindIII/SmaI LEU2 containing fragment from pJJ283 (Jones and Prakash 1990), to form pT7Blue YKU70::LEU2. In order to obtain YKU70 disruptants Leu+ colonies were selected after transformation of YPH250 with a 2884 bp NdeI/SmaI fragment from pT7BlueYKU70::LEU2. The ExpandTM High Fidelity System (Boehringer Mannheim) was used 15 according to the supplied protocol to amplify a 3285 bp fragment from the LIG4 locus with primers dnl4p1 5'-CGTAAGATTCGCCGAGTATAG-3' and dnl4p2 5'-CGTTTCAAATGGGACCACAGC-3'. The PCR product was cloned into pGEMT (Promega), resulting in pGEMTLIG4. A 1326 bp BamHI/XhoI 20 fragment from pJJ215 (Jones and Prakash 1990) containing the HIS3 gene was inserted into the BamHI and XhoI sites of pIC20R, resulting in pIC20RHIS3. A 782 bp EcoRI fragment from the LIG4 ORF was replaced with a 1367 bp EcoRI HIS3 containing fragment from pIC20RHIS3 to construct 25pGEMTLIG4::HIS3. In order to obtain LIG4 disruptants His+ colonies were selected after transformation of YPH250 with a 3854 bp NcoI/NotI fragment from pGEMTLIG4::HIS3. In order to obtain RAD50 disruptants YPH250 was transformed with a EcoRI/BglII fragment from pNKY83 and Ura+ colonies were selected (Alani et al 1989). A rad50::hisG strain was obtained by

selecting Ura colonies on selective medium containing 5-FOA. Similarly

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RAD51 disruptants were obtained were obtained after transformation of YPH250 with a RAD51::LEU2 XbaI/PstI fragment from pDG152 and selection of Leu+ colonies (Schiestl et al 1994). The TRP1 marker in pSM21 (Schild et al. 1983) was replaced with a BglII/XbaI LEU2 containing fragment from pJJ283 (Jones and Prakash, 1990). This resulted in pSM21LEU2. Leu+RAD52 disruptant colonies were selected after transformation of YPH250 with the RAD52::LEU2 BamHI fragment from pSM21LEU2. Disruption constructs were transformed to YPH250 by the lithium acetate transformation method as described (Gietz et al. 1992; Schiestl et al. 1993). Disruption of YKU70, LIG4, RAD50, RAD51 and RAD52 was confirmed by PCR and Southern blot analysis.

Table 1: Yeast strains

Table 1. Teast s			
Strain	Genotype	Reference	
YPH250	MATa, ura3-52, lys2-801, ade2-101,	(Sikorski and	
	trp1-∆1, his3-∆200, leu2-∆1	Hieter 1989)	
YPH250rad51	MATlpha, $ura 3-52$, $lys 2-801$, $ade 2-101$,	This study	
	trp1-∆1, his3-∆200, leu2-∆1,	-	
	rad 51::LEU2		
${\rm YPH}250 rad 52$	MATα, ura3-52, lys2-801, ade2-101,	This study	
	trp1-∆1, his3-∆200, leu2-∆1,		
	rad 52::LEU2		
YPH250yku70	MATa, ura3-52, lys2-801, ade2-101,	This study	
	trp1-∆1, his3-∆200, leu2-∆1,		
	yku70::LEU2		
YPH250rad50	MATα, ura3-52, lys2-801, ade2-101,	This study	
	trp1-∆1, his3-∆200, leu2-∆1,rad50::hisG		
YPH250lig4	MATα, ura3-52, lys2-801, ade2-101,	This study	
	trp1-∆1, his3-∆200, leu2-∆1, lig4::HIS3		
JKM115	Δho, Δhml::ADE1, MATα, Δhmr::ADE1,	(Moore and	
	ade1, $leu2$ -3,112, $lys5$, $trp1$:: $hisG$,	Haber 1996)	
	ura 3-52		
JKM129	Δho, Δhml::ADE1, MATα, Δhmr::ADE1,	(Moore and	
	ade1, leu2-3,112, lys5, trp1::hisG,	Haber 1996)	
	ura3-52, xrs2::LEU2		
JKM138	Δho, Δhml::ADE1, MATα, Δhmr::ADE1,	(Moore and	
1	ade1, leu2-3,112, lys5, trp1::hisG,	Haber 1996)	
	ura3-52, mre11::hisG	,	
YSL204	Δho, HMLα, MATα, HMRα, ade1-100,	(Lee et al.	
	leu2-3,112, lys5, trp1::hisG, ura3-52,	1999)	
	hisG'-URA3-hisG', sir4::HIS3	-	

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Construction of binary vectors.

To construct pSDM8000 a 1513 bp PvuII/EcoRV fragment carrying the KanMX marker was obtained from pFA6a (Wach et al. 1994) and was ligated into the unique HpaI site of pSDM14 (Offringa 1992). pSDM8001 was made in three cloning steps. A 1476 bp BamHI/EcoRI fragment carrying the KanMX marker was obtained from pFA6a and ligated into BamHI and EcoRI digested pIC20H to form pIC20HkanMX. The KanMX marker was inserted between the PDAI flanks by replacement of a 2610 bp BgIII fragment from pUC4E1α10 (Steensma et al. 1990) with a 1465 BgIII fragment from pIC20HkanMX. A 3721 bp XhoI/KpnI fragment from this construct was inserted into the XhoI and KpnI sites of pSDM14. The binary vectors pSDM8000 and pSDM8001 were introduced into Agrobacterium tumefaciens LBA1119 by electroporation (den Dulk-Ras and Hooykaas 1995).

15 Cocultivations / T-DNA transfer experiments.

Cocultivations were performed as described earlier with slight modifications (Bundock et al. 1995). Agrobacterium was grown overnight in LC medium. The mix of Agrobacterium and S. cerevisiae cells was incubated for 9 days at 20°C. G418 resistant S.cerevisiae strains were selected at 30°C on YPAD medium containing geneticin (200 µg/ml) (Life Technologies/Gibco BRL).

Vectorette PCR.

Chromosomal DNA was isolated using Qiagen's Genomic Tips G/20 per manufacters protocol. 1-2 µg of Genomic DNA was digested with EcoRI, ClaI, PstI or HindIII and run on a 1% TBE-gel. Non-radioactive Southern blotting was performed. The membrane was hybridized with a digoxigenine-labeled kanMX probe to determine the size of T-DNA/genomic DNA fragments (EcoRI and ClaI for RB containing fragments and PstI and HindIII for LB containing fragments). The kanMX probe, a 792 bp internal fragment of the KanMX

marker, was made by PCR using primers kanmxp1 5'-AGACTCACGTTTCGAGGCC-3' and kanmxp2 5'-

TCACCGAGGCAGTTCCATAG-3' and a Non-Radioactive DNA Labeling and Detection kit (Boehringer Mannheim). The enzyme showing the smallest band on blot was used for Vectorette PCR, in order to amplify the smallest junction sequence of T-DNA and genomic DNA. Vectorette PCR was performed as described

(http://genomewww.stanford.edu/group/botlab/protocols/vectorette.html). The ExpandTM High Fidelity System (Boehringer Mannheim) was used to amplify fragments larger than 2.5 kb, whereas sTaq DNA polymerase (SphaeroQ) was used for amplification of fragments smaller than 2.5 kb. Primer kanmxp3 5'-TCGCAGGTCTGCAGCGAGGAGC-3' and kanmxp4 5'-

TCGCCTCGACATCATCTGCCCAG-3' were used to amplify RB/genomic DNA and LB/genomic DNA junction sequences, respectively.

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T7 DNA Polymerase sequencing.

Vectorette PCR products were cloned in pGEMTEasy (Promega) and sequenced using the T7 polymerase sequencing kit (Pharmacia) according to manufacturers protocol. In order to obtain sequences flanking the RB and LB, primers kanmxp5 5'-TCACATCATGCCCCTGAGCTGC-3' and kanmxp4 were used, respectively.

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RESULTS

1. Binary vectors for T-DNA transfer to yeast.

It was previously demonstrated that Agrobacterium tumefaciens is able to transfer its T-DNA not only to plants but also to another eukaryote, namely the yeast Saccharomyces cerevisiae (Bundock et al. 1995). T-DNA carrying homology with the yeast genome was shown to become integrated by homologous recombination. T-DNA lacking any homology with the yeast genome was integrated randomly into the genome by IR like in plants (Bundock et al. 1995, (Bundock and Hooykaas 1996). The T-DNA used in these experiments carried the S.cerevisiae URA3 gene for selection of Ura+ colonies after T-DNA transfer to the haploid yeast strain RSY12(URA34). However, in this system only yeast strains could be used in which the URA3 gene had been deleted to avoid homology between the incoming T-DNA and the S.cerevisiae genome.

We wanted to setup a system in which T-DNA transfer to any yeast strain could be studied. Therefore, two new binary vectors were constructed using the dominant marker kanMX (Wach et al 1994)which confers resistance against geneticin (G418). The T-DNA of pSDM8000 carries only the KanMX marker. Since this KanMX marker consists of heterologous DNA, lacking any homology with the S.cerevisiae genome, we would expect this T-DNA to integrate by IR at a random position in the yeast genome. To be able to compare this with T-DNA integration by homologous recombination pSDM8001 was constructed. The T-DNA of pSDM8001 carries the KanMX marker flanked by sequences from the S.cerevisiae PDA1 locus. The PDA1 sequences have been shown to mediate the integration of T-DNA by HR at the PDA1 locus on chromosome V (Bundock et al. 1995).

Cocultivations between Agrobacterium strains carrying pSDM8000 and pSDM8001, respectively, and the haploid yeast strains YPH250 and JKM115, respectively, were carried out as described in the experimental part. G418

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resistant colonies were obtained at low frequencies for YPH250 (1.6 x 10^{-7}) and JKM115 (1.2 x 10^{-5}) after T-DNA transfer from pSDM8000 (Table 2). T-DNA transfer from pSDM8001 generated G418 resistant colonies at higher frequencies (2.4 x 10^{-5} for YPH250 and 1.8 x 10^{-4} JKM115, Table 2). The ratio of homologous recombination versus illegitimate recombination is determined by comparing the frequencies of G418 resistant colonies obtained from cocultivations using either pSDM8001 or pSDM8000. This showed that a T-DNA from pSDM8001 was 150-fold more likely to integrate than a T-DNA from pSDM8000 in YPH250 (Table 2). A similar difference was previously seen using T-DNAs with the URA3 marker (Bundock and Hooykaas 1996). In contrast, T-DNA from pSDM8001 was only 16-fold more likely to integrate than a T-DNA from pSDM8000 in JKM115. There was no significant difference in the frequency of T-DNA transfer to these two yeast strains as was determined by T-DNA transfer experiments in which a T-DNA, that carried the KanMX marker and the yeast 2 micron replicon, was employed. Therefore, the differences in the frequencies of T-DNA integration by HR and IR between the yeast strains YPH250 and JKM115, respectively, is most likely contributed to differences in the capacities of their HR and IR recombination machineries.

We confirmed by PCR that the T-DNA from pSDM8001 became integrated at the *PDA1* locus by homologous recombination (data not shown). In order to find out whether the T-DNA from pSDM8000 had integrated randomly by IR yeast target sites for integration were determined from 8 G418 resistant YPH250 colonies by Vectorette PCR (for detailed description see materials and methods). Chromosomal DNA was isolated and digested with a restriction enzyme that cuts within the T-DNA. A Vectorette was ligated to the digested DNA and a PCR was performed using a T-DNA specific and a Vectorette specific primer. The PCR product obtained was cloned into pGEMTEasy and sequenced using a T-DNA specific primer. The position of the T-DNA insertion was determined by basic BLAST search of the yeast genome (http://www-genome.stanford.edu/SGD). We were thus able to map the position

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of the T-DNA insertions of all 8 G418 resistant colonies analyzed. They were present at different positions spread out over the genome. Comparison of the T-DNA sequence and yeast target site sequences did not reveal any obvious homology. These data show that the T-DNA from pSDM8000 had integrated via an IR mechanism as expected.

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The following characteristics have previously been observed for T-DNAs integrated by IR: a) the 3' end of the T-DNA is usually less conserved compared to the 5' end, b) microhomology is sometimes present between the T-DNA ends and the target site, c) integration is often accompanied by small deletions of the target site DNA (Matsumotot et al. 1990), (Gheysen et al. 1991, (Mayerhofer et al. 1991), (Bundock and Hooykaas 1996). Similar characteristics were seen in the currently analyzed 8 T-DNA insertions. In 3 strains we observed microhomology of 2 – 6 bp between the LB and yeast target site (figure 1, WT.51 was taken as an example). In 5 strains deletions of 1 – 5 bp of yeast target site DNA was found and we observed deletions, varying from 1 – 112 bp, of the 3' end of the T-DNA in 7 out of 8 analyzed strains. In only 1 strain the 3' end appeared to be intact. The 5' end of the T-DNA was conserved in almost all strains. In only 2 strains we observed small deletions of 1 and 2 bp at the 5' end of the T-DNA.

Thus, we can conclude that the T-DNA from pSDM8000 had integrated via the same IR mechanism described before.

Table 2: Frequencies of T-DNA integration by IR relative to integration by HR in recombination defective yeast strains

Strain	Genotype	Freq. of	Freq. of	Absolute	Standardized
		IR^a	HR	IR/HR	IR/HR ratio ^c
				$ratio^b$	
YPH250	WT	1.6 x 10 ⁻⁷	2.4 x 10 ⁻⁵	0.007	1
YPH250	$rad51$ $_{\Delta}$	1.4×10^{-7}	1.5 x 10 ⁻⁶	0.09	14
rad51					
YPH250	rad52s	3.8×10^{-7}	2.5×10^{-6}	0.15	23
rad52					
YPH250	. yku70∆	<3.2 x 10 ⁻⁹	3.3×10^{-5}	< 0.0001	<0.01
yku 70					
YPH250	$rad50$ $_{\Delta}$	8.0×10^{-9}	3.5×10^{-5}	0.0002	0.03
rad50					
YPH250	lig4⊿	3.7×10^{-9}	2.3×10^{-5}	0.0002	0.02
lig4					
JKM115	WT	1.2×10^{-5}	1.8 x 10 ⁻⁴	0.07	1
JKM129	xrs24	2.7×10^{-7}	5.1×10^{-5}	0.005	0.08
JKM138	$mre11\Delta$	2.9×10^{-7}	7.5×10^{-5}	0.004	0.06
YSL204	sir4∆	1.5×10^{-7}	1.8 x 10 ⁻⁵	0.008	0.13

<sup>a Averages of 2 or more independant experiments are shown. Frequencies are
depicted as the number of G418 resistant colonies devided by the output number of yeast cells (cells/ml).</sup>

^b The frequency of T-DNA integration by IR (pSDM8000) devided by the frequency of T-DNA integration by HR (pSDM8001).

^c The ratio of IR/HR in the mutant strain devided by the ratio of IR/HR in the wildtype strain.

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2. Host-specific factors involved in random T-DNA integration.

The observation that the T-DNA from pSDM8000 integrates by IR into the yeast genome allowed us to use this system to study the effect of host factors on the process of integration. Many proteins involved in various forms of DNA recombination have been identified in yeast. In order to determine the roles of a representative set of these enzymes in T-DNA integration, we compared T-DNA transfer and integration in wildtype yeasts with that of strains carrying disruptions of the genes encoding several recombination proteins. The RAD51, RAD52, KU70, RAD50 and LIG4 genes were deleted from YPH250 using the one step disruption method. Yeast strains carrying deletions in MRE11, XRS2 and SIR4 in the JKM115 background were kindly provided by Dr. J. Haber. The results of cocultivations with these yeast strains are shown in Table 2.

In rad51 and rad52 mutants, which are impaired in homologous recombination, the frequency of T-DNA integration by HR was 16- and 9-fold lower, respectively, than observed for the wildtype (Table 2). This showed that RAD51 and RAD52 play a role in T-DNA integration by homologous recombination. In the IR defective ku70, rad50, lig4, mre11, xrs2 and sir4 mutants the frequency of T-DNA integration by HR did not differ significantly from that observed for wildtype (Table 2). This showed that these genes do not play a role in T-DNA integration by homologous recombination.

The frequency of T-DNA integration by IR in a rad51 mutant did not differ significantly from that observed for wildtype, whereas in a rad52 mutant the frequency was about 2-fold higher (Table 2). This showed that RAD51 and RAD52 are not involved in T-DNA integration by IR. The product of the RAD52 gene may compete with IR-enzymes for the T-DNA and thereby inhibits integration by IR to some extent. Strikingly, in rad50, mre11, xrs2, lig4 and sir4 mutants the frequency of T-DNA integration by IR was reduced dramatically: 20- to more than 40-fold (Table 2). T-DNA integration by IR seemed to be completely abolished in the ku70 mutant. We did not obtain any

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G418 resistant colonies from several cocultivation experiments. This strongly suggests that KU70 plays an important role in random T-DNA integration in yeast.

Since T-DNA integration by HR is normal in these mutants, these results clearly show that the yeast genes KU70, RAD50, MRE11, XRS2, LIG4 and SIR4 are involved in T-DNA integration by illegitimate recombination.

3. Chromosomal distribution of integrated T-DNA copies in IR defective S.cerevisiae.

From several cocultivation experiments with the rad50, mre11, xrs2, lig4 and sir4 mutants we obtained a small number of G418 resistant colonies. The T-DNA structure was determined for a number of these lines. To this end chromosomal DNA was isolated from these G418 resistant colonies and subjected to vectorette PCR to amplify junction sequences of genomic DNA and T-DNA. PCR products were cloned and sequenced. The yeast sequences linked to the T-DNA were used in a BLAST search at http://www-genome.stanford.edu/SGD to determine the T-DNA integration sites.

Strikingly, analysis of LB/genomic DNA junctions revealed that in 2 out of 3 rad50, 4 out of 6 mre11 and 2 xrs2 strains analyzed, T-DNAs had integrated in telomeres or subtelomeric regions (rad50k.1, rad50k.6, mre11k.8, mre11k.11, mre11k.14, mre11k.17, xrs2k.1 and xrs2k.17; Table 3 and figure 1). S. cerevisiae telomeres generally consist of one or more copies of the Y element followed by telomerase-generated C(1-3)A/TG(1-3) repeats (Zakian 1996). In 2 rad50 strains, 2 mre11 strains and 1 xrs2 strain the LB was found to be fused to this typical telomerase-generated C(1-3)A/TG(1-3) repeat (rad50k.1, rad50k.6, mre11k.14, mre11k.17 and xrs2k.1; figure 1). Besides, we also found one T-DNA insertion in a Ty LTR element in the mre11 mutant and 2 insertions in the rDNA region, present in chromosome XII, in the mre11 and rad50 mutants (mre11k.5, mre11k.4 and rad50k.5, respectively; Table 3 and figure 1).

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The 3' end of the T-DNA was truncated in all strains. Deletions of 3 – 11 bp of the 3'end of the T-DNA were observed (figure 1). Microhomology between the 3' end of the T-DNA and yeast target site was only found in 2 lines (5 bp in mre11k.4 and 4 bp in mre11k.14; figure 1). For the T-DNA copies present at the yeast telomeres, the RB/genomic DNA junction sequences could not be obtained from these strains using vectorette PCR. This was only possible for the rad50 and mre11 strains carrying the T-DNA in the rDNA region on chromosome XII. In both strains the RB was intact and no homology between the 5' end of the T-DNA and the yeast target site was found (data not shown in figure 1).

Previously, target sites for T-DNA integration in the genome of S.cerevisiae strain RSY12 were determined (Bundock and Hooykaas 1996), (Bundock 1999. In 4 out of 44 strains analyzed, T-DNA copies were integrated in rDNA, Ty LTR elements (in 2 strains) and a subtelomeric located Y element, respectively. In addition, we determined the position of T-DNA integration in ten S.cerevisiae YPH250 strains. We did not find any T-DNA insertions in rDNA, LTR elements or subtelomeric/telomeric regions amongst these ten. Pooling all insertions analyzed in wildtype (54), in 2 out of 54 strains analyzed (4%) insertions were found in a Ty LTR element and in two other strains in the rDNA repeat (2%) and a subtelomeric region (2%), respectively. In contrast, we report here that T-DNA in yeast strains mutated in RAD50, MRE11 or XRS2 T-DNA integrates preferentially in (sub)telomeric regions (8 out of 11 lines: ~73%) of rad50, mre11 and xrs2 mutants (table 3). From the remaining strains two T-DNAs were present in rDNA and one in a Ty LTR element, respectively. Apparently, the rDNA repeat is also a preferred integration site in these mutants (~18% vs. ~2% in the wildtype).

Telomeres consist of a large array of telomerase-generated C(1-3)A/TG(1-3) repeats (~350 bp). In the subtelomeric regions two common classes of Y elements, 6.7 and 5.2 kb, can be found (in most strains chromosome I does not contain Y) (Zakian and Blanton 1988), making the average size of these

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regions ~6,0 kb. Thus, the yeast genome contains $(16 \times 2 \times 0.35) + (15 \times 2 \times 6.0) = 191$ kb of subtelomeric/telomeric sequences. The yeast genome is 12,052 kb in size, which means that only 1.6% of the genome consists of subtelomeric/telomeric sequences. In accordance with this, we observed in only 2% of the wildtype strains T-DNA copies inserted in a subtelomeric region, which we would expect on the basis of random T-DNA integration. In contrast, in the rad50, mre11 and xrs2 mutants 73% of the T-DNA insertions were found in the (sub)telomeric region.

Analysis of 7 lines revealed that in the *sir4* mutant T-DNA was integrated randomly into the yeast genome. So, although *SIR4* has an effect on the efficiency of T-DNA integration by IR, the pattern of T-DNA distribution in the transformants seems similar as in the wildtype strain. In the *sir4* mutant T-DNA integration by IR was characterized by truncation of the 3' end of the T-DNA, deletions at the target site and microhomology between the LB and the target site (data not shown), like this was observed for T-DNA integration by IR in the wildtype.

These results clearly show that in the rad50, mre11 and xrs2 mutants the T-DNA, if integrated at all, becomes preferentially inserted in telomeres or subtelomeric regions and that the genomic distribution of integrated T-DNAs is altered when compared to wildtype. However, disruption of SIR4 did affect the efficiency of T-DNA integration by IR, but not the characteristics of this process.

Table 3: genomic distribution of T-DNA integrated by IR in rad50, mre11 and xrs2 mutants in comparison with the wildtype after T-DNA transfer from $^-$ pSDM8000

Yeast strain	(Sub)Telomeric region	LTR	rDNA	Elsewhere
rad50 mutant	2	0	1	0
mre11 mutant	4	1	1	0
xrs2 mutant	2	0	0	0
wildtype	1	2	1	50

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DESCRIPTION OF FIGURES

Figure 1: Junction sequences of T-DNA and S.cerevisae genomic DNA.

S.cerevisiae YPH250 (WT), rad50, mre11 and xrs2 strains were cocultivated with LBA1119(pSDM8000). G418 resistant colonies were obtained.

Chromosomal DNA was isolated and subjected to Vectorette PCR to determine the sequence of genomic DNA flanking the T-DNA. Position of T-DNA integration was determined by basic BLAST search of the yeast genome at http://www.genome-stanford.edu/SGD. The Watson strand of genomic DNA that is fused to the LB or RB is shown in italics. Bold sequences represent sequence homology between the LB and target site. The filler DNA sequence is underlined and depicted in italics. The numbers above the LB sequences represents the number of bp deleted from the LB. Tel. = telomeric, Subtel. = subtelomeric and Int. = intergenic.

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Figure 2: Alignment of KU70 homologues. Sc = Saccharomyces cerevisiae, Hs = Homo sapiens and At = Arabidopsis thaliana. Perfect identity is depicted as black boxes, homology is depicted as grey boxes and dashes were used to optimise alignment.

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Figure 3: Alignment of LIG4 homologues. Sc = Saccharomyces cerevisiae, Hs = Homo sapiens and At = Arabidopsis thaliana. Perfect identity is depicted as black boxes, homology is depicted as grey boxes and dashes were used to optimise alignment.

Figure 4: Alignment of MRE11 homologues. Sc = Saccharomyces cerevisiae, Hs = Homo sapiens and At = Arabidopsis thaliana. Perfect identity is depicted as black boxes, homology is depicted as grey boxes and dashes were used to optimise alignment.

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Figure 5: Alignment of RAD50 homologues. Sc = Saccharomyces cerevisiae, Hs = Homo sapiens and At = Arabidopsis thaliana. Perfect identity is depicted as black boxes, homology is depicted as grey boxes and dashes were used to optimise alignment.

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Figure 6: Alignment of XRCC4 homologues. Sc = Saccharomyces cerevisiae, Hs = $Homo\ sapiens\ and\ At = Arabidopsis\ thaliana$.

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Claims

- 1. A method to direct integration of a nucleic acid of interest to a predetermined site, whereby said nucleic acid has homology at or around the said pre-determined site, in a eukaryote with a preference for non-homologous recombination, comprising steering an integration pathway towards homologous recombination.
- 2. A method to direct nucleic acid integration according to claim 1, comprising providing a mutant of a component involved in non-homologous recombination.
- 3. A method to direct nucleic acid integration according to claim 1 or 2, comprising inhibiting a component involved in non-homologous recombination.
- 4. A method according to claim 2 or 3 wherein said component involved in non-homologous recombination comprises ku70, rad50, mre11, xrs2, lig4 or sir4.
- 5. A method to direct integration of a nucleic acid of interest to a predetermined site according to anyone of claims 1 to 3, wherein said nucleic acid sequence of interest is essentially replacing a sequence within said eukaryote.
- 6. A method to direct integration of a nucleic acid of interest to a pre-20 determined site according to claim 5, wherein said component involved in non-homolgous recombination comprises rad50 or xrs2.
 - 7. A method to direct integration of a nucleic acid of interest to a subtelomeric and/or telomeric region in a eukaryote with a preference for non-homologous recombination by providing a mutant of a component involved in non-homologous recombination.
 - 8. A method to direct integration of a nucleic acid of interest to a subtelomeric and/or telomeric region in a eukaryote with a preference for nonhomologous recombination, comprising inhibiting a component involved in non-homologous recombination

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- 9. A method to direct integration according to claim 7 or 8 wherein said component involved in non-homologous recombination comprises rad50, mre11 or xrs2.
- 10. A method according to anyone of claims 1 to 9 wherein said eukaryote comprises yeast, a fungus or an animal.
- 11. A method according to anyone of claims 1 to 10, wherein said nucleic acid of interest is delivered to a cell of said eukaryote by *Agrobacterium*.
- 12. A method according to anyone of claims 1-11 comprising transiently inhibiting integration via non-homologous recombination.
- 10 13. A method according claim 12 wherein said transiently inhibiting is provided by an *Agrobacterium* Vir-fusion protein capable of inhibiting a component involved in non-homologous recombination.
 - 14. A method to direct nucleic acid integration according to claim 13 wherein said *Agrobacterium* Vir fusion protein comprises VirF or VirE2.
- 15 14. A method according to claim 13 or 14 wherein said component involved in non-homologous recombination comprises ku70, rad50, mre11, xrs2, lig4 or sir4.
 - 15. A method according to anyone of the aforegoing claims wherein said nucleic acid of interest comprises an inactive gene to replace an active gene.
- 20 16. A method according to anyone of claims 1-14, wherein said nucleic acid of interest comprises an active gene to replace an inactive gene.
 - 17. A method according to anyone of claims 1-14, wherein said nucleic acid of interest encodes a therapeutic proteinaceous substance.
- 18. A method according to anyone of claims 1-14, wherein said nucleic acid of interest encodes a substance conferring resistance for an antibiotic substance to a cell.
 - 19. A method according to anyone of claims 1-14, wherein said nucleic acid of interest confers a desired property to said eukaryotic cell.
- 20. A method according to anyone of the aforegoing claims wherein said nucleic acid of interest is part of a gene delivery vehicle.

21. Use of a method according to anyone of claims 1 to 20 for improvement of gene targeting efficiency.

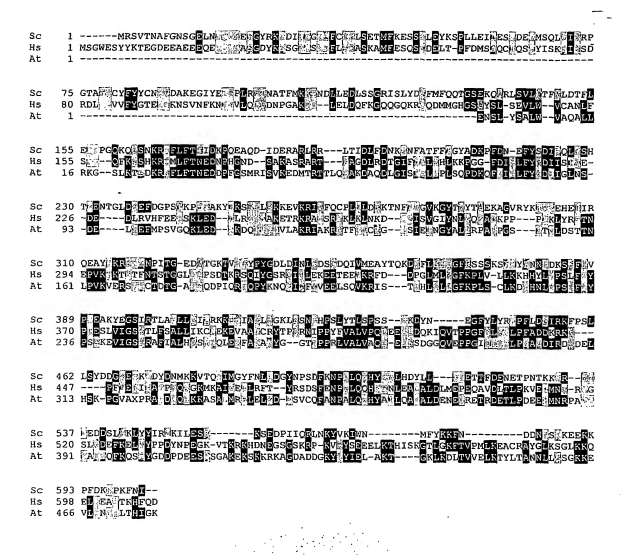
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Fig. 1

Strain	LB' T-DNA RB' CAGGATATATTCAATTGTAAAT-CTCCGA-GG C	hromosome, coordinate and location
WT.51	-4 5'ATTGTATTATATTCAATTGTAAAT-CTCCGA-GGTA 3'	XIV, 185311 (1 bp of target site DNA deleted), int. region
rad50k.1	5'TGTGGGTGTGATATTCAATTGTAAAT-CTC—CGA-GG 3'	XV, 1091277, tel. region
rad50k.5	-7 5' <i>GGGGCATCAG</i> TATTCAATTGTAAAT-CTCCGA-GG 3'	XII, 465986, rDNA region
rad50k.6	-25 5' GAGGTAGATGTGAGAGAGTGTGTGTGGGTGTGAAGTCGA 3'	XV, 1091276, tel. region
	-3	
mre11k.4	5' TCTGGTAGATATATTCAATTGTAAAT-CTCCGA-GG 3'	XII, 459692/468829, rDNA region
mre11k.5	5'CACATATTCTCATTCAATTGTAAAT-CTCCGA-GG 3'	VII/X/XIII, 536090 OR 541678/ 472487 OR 483659/196667, LTR
mre11k.8	5' CGACTACTTTATATCCAATTGTAAAT-CTCCGA-GG 3'	XIV, 6060, subtel. region
mre11k.11	5' GAAGAACCCATTATTCAATTGTAAAT-CTCCGA-GG 3'	XIV, 4866, subtel. region
mre11k.14	5' TGGGTGTGGGTTATTCAATTGTAAAT-CTCCGA-GG 3'	VIII, 562588, tel. region
mre11k.17	5' TGGGTGTGTGTTCAATTGTAAAT-CTCCGA-GG 3'	XII, 5727, subtel. region
	-10	
xrs2k.1	5' TGTGTGGGTGTGGGTCAATTGTAAAT-CTCCGA-GG 3'	IX/X, 69/52, tel. region
xrs2k.17	-1 5' CGTCAAGGATATATTCAATTGTAAAT-CTCCGA-GG 3'	XII, 1071797, subtel. region

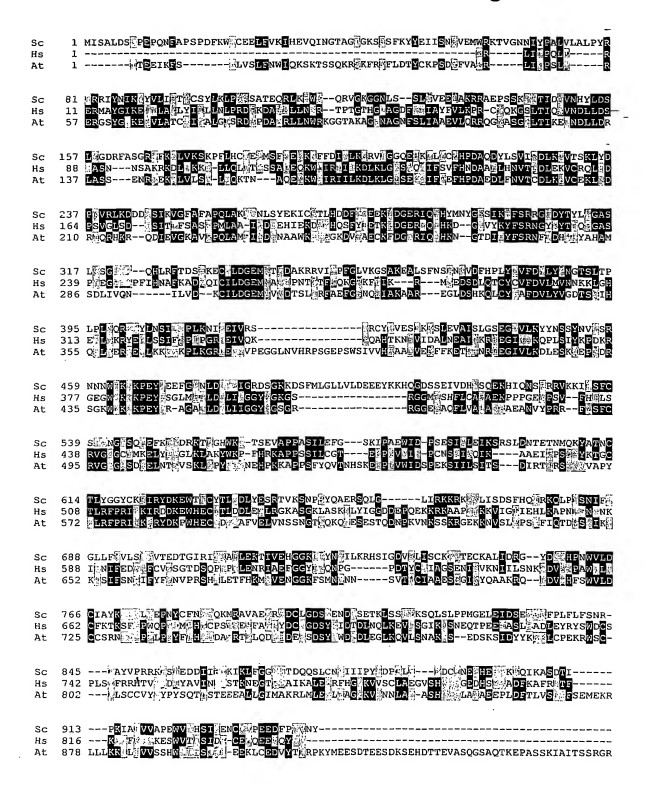
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Fig. 2



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Fig. 3



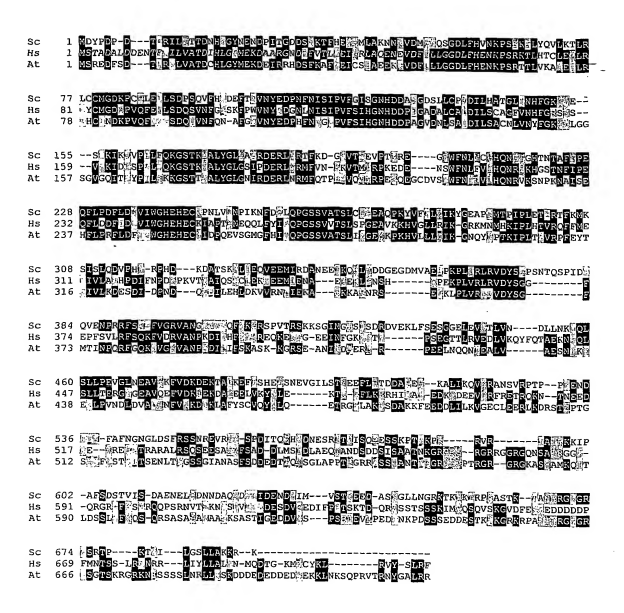
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Fig. 3, Contd.

SC Hs		SNTRAVKRGRSSTNSLQRVQRRRGKQPSKISGDETEESDASEEKVSTRLSDIAEETDSFGEAQRNSSRGKCAKRGKSRVG
Sc Hs		
	1037	${\tt QTQRVQRSRRGKKAAKIGGDESDENDELDGNNNVSADAEEGNAAGRSVENEETREPDIAKYTESQQRDNTVAVEEALQDS} \\$
Sc		
Нs		
At	1117	${\tt RNAKTEMDMKEKLQIHEDPLQAMLMKMFPIPSQKTTETSNRTTGEYRKANVSGECESSEKRKLDAETDNTSVNAGAESDV}$
Sc		
Нs		
Αt	1197	VPPLVKKKKVSYRDVAGELLKDW

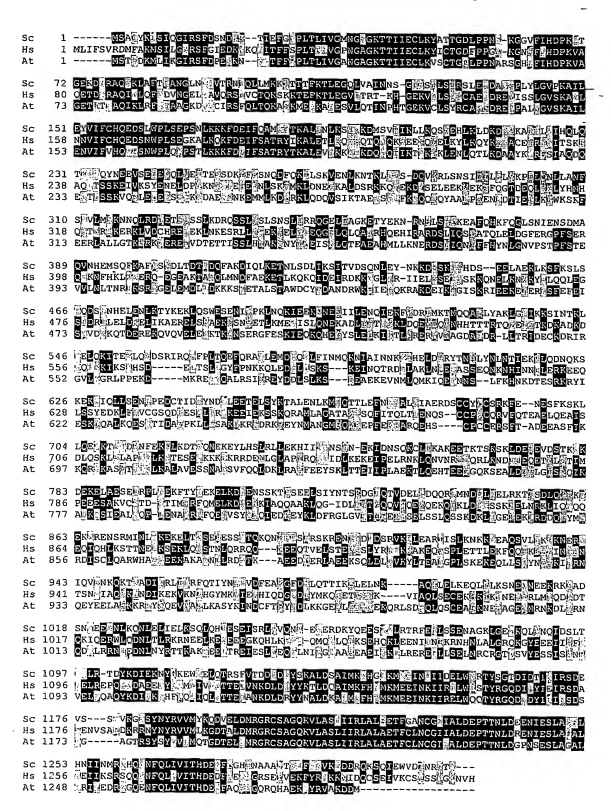
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Fig. 4



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Fig. 5



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Fig. 6

	1	50
At	(1)	MIGVDSKSSSTTFIETMVESEKTKHTC
Hs	(1)	MERKISR
Sc	(1)	MSQLTEFISCIPVVNEEQNEEDERGLCKIQIEDGAMLETLDENSLSGLRI
Consensus	(1)	IETL E SKIS
		51
At ((28)	LRLEISGADPIFVKGTWHNSRFDISVTDGSSS
Hs	(8)	
Sc ((51)	EKMLYSEGTGIFSKSSFGINDLRIFTGENIDEESKKYVWYELLKMLTGHK
Consensus ((51)	IKLLIS A IFLK SW S F ISLTDG SA
		101
At ((60)	101 150WICNAŢEËËVÄERAAQWĎQPVSEYĹ-KLAEQYĽGFQ
•	(43)	WIGTVSESEISQEADDMAMEKGKYVGELRKALLSGA
	(01)	VYIASLDEKVVFTKWTCRMQDDEVWKVVMELESSAIIRKIAELTLHPV
· · · · · · · · · · · · · · · · · · ·	101)	WTC MSEDEVA A DLD S YI KLAE L
.		151 200
	(95)	
Hs (Sc (1	(19)	GPADVYTFNFSKESCYFFFEKNLKDVSFRLGSFNLEKVENPAEVIRELIC KKGEIDLFEMADKLYKDICCVNDSYRNIKESDSSNRNRVEQLARERELLD
	151)	
	•	
		201 250
At (1	L45)	FLMEANIRLSEEVUNKTR-SFEKMRSEAERCLAQGEKLCDEKTEFESATY
		YCLDTIAENQAKNEHLQK-ENERLLRDWNDVQGRFEKCVSAKEALETDLY
	199) 201)	
Consensus (2	2017	FLLET E S MM K E EKLIRE DILAQ EK LSDKD LESALY
		251 300
•	194)	
	178)	
	249) 251)	NTEVOKPISELNSPGKRMKRRKTVVEPONLOKKLKDTSRRRANRKISNOS KFISVLNA KK KIR AL KL A RE
Consensus (2	231)	KFISVLNA KK KIR AL KL A RE
		301 350
- •	223)	EESTDKAESFESGRSDDEKSEEEASKKATSSKARGGKRAARS
	215)	
		VIKMEDDDFDDFQFFGLSKRPIITAKDKLSEKYDDITSFGDDTQSISFES
Consensus (3	301)	EDS DK FD SK DD A DDLS KASAA SKGD S S
		351 400
At (2	265)	700
Hs (2	265)	
	349)	
Consensus (3	351)	DAK GAD D DS SAS
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